

Impact of surface emissions to the zonal variability of tropical tropospheric ozone and carbon monoxide for November 2004

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Abstract

The chemical and dynamical processes governing the zonal variability of tropical tropospheric ozone and carbon monoxide are investigated for November 2004 using satellite observations, *in-situ* measurements, and chemical transport models in conjunction with inverse estimated surface emissions. Vertical ozone profile estimates from the Tropospheric Emission Spectrometer (TES) and ozone sonde measurements from the Southern Hemisphere Additional Ozonesondes (SHADOZ) network show the so-called zonal “wave-one” pattern, which is characterized by peak ozone concentrations (70-80 ppb) centered over the Atlantic, as well as elevated concentrations of ozone over Indonesia and Australia (60-70 ppb) in the lower troposphere. Observational evidence from TES CO vertical profiles, Ozone Monitoring Instrument (OMI) NO₂ columns point to regional surface emissions as an important contributor to the elevated ozone over Indonesia. This contribution is investigated with the GEOS-Chem chemistry and transport model using surface emission estimates derived from an optimal inverse model, which was constrained by TES and Measurements Of Pollution In The Troposphere (MOPITT) CO profiles (Jones et al., 2007). These *a posteriori* estimates, which were over a factor of 2 greater than climatological emissions, reduced differences between GEOS-Chem and TES ozone observations by 30-40% and led to changes in GEOS-Chem upper tropospheric ozone of up to 40% over Indonesia. The remaining residual differences can be explained in part by upper tropospheric ozone produced from lightning NO_x in the South Atlantic. Furthermore, model simulations from GEOS-Chem indicate that ozone over Indonesian/Australian is more sensitive to changes in surface emissions of NO_x than ozone over the tropical Atlantic.

1 Introduction

The distribution of tropical tropospheric ozone is governed by the complex interplay of chemistry and dynamics. Ozone can be generated from surface emissions such as biomass burning, forest fires and fossil fuels through the production of carbon monoxide and hydrocarbons in the presence of nitrogen oxides (NO_x) (Jacob et al., 1996). The monthly distribution and

intensity of these emissions can vary between South America, sub-equatorial Africa, and Indonesia/Australia (Arellano et al., 2006; Duncan et al., 2003b,a). Furthermore, the production and distribution of ozone from these emissions depends nonlinearly on the type of emission, the intensity of those emissions, and prevailing meteorological conditions. In the middle and upper troposphere, ozone can be generated efficiently through lightning-based production of NO_x (Pickering et al., 1998; Martin et al., 2000, 2002; Jenkins and Ryu, 2004b,a). Tropospheric ozone can be transported globally where it can impact the oxidative capacity of the global atmosphere, radiative forcing of the climate system, and air quality (Fishman et al., 1979, 1991; Fishman and Larsen, 1987; Lacis et al., 1990; Kiehl et al., 1999; Portmann et al., 1997; Naik et al., 2005; Jacob, 1999; Li et al., 2002).

Earth-observing satellites provide a rich suite of data to investigate the processes controlling tropical tropospheric ozone. In particular, the Tropospheric Emission Spectrometer (TES), aboard NASA's Aura spacecraft, adds a unique observational dataset that includes vertical estimates of both ozone and a key signature of pollution, carbon monoxide. Co-located measurements of ozone and CO can help distinguish between natural and anthropogenic sources of ozone (Zhang et al., 2006) and the vertical information can aid in disentangling the meteorological processes driving the redistribution of ozone (Jourdain et al., 2007). This information will be crucial to unravel the impact of surface emissions on free tropospheric ozone.

We investigate the impact of surface emissions on the distribution of ozone in the tropical troposphere based on an integrated approach that combines multiple satellite data, sonde measurements, chemistry and transport modeling under the framework of data assimilation and linear optimal estimation. Satellite observations will provide insight into the sources of ozone precursors, as well as the distribution of precursors and concomitant ozone. The analysis is focused over the southern hemisphere during November 2004, which marks a transitional period between Austral winter and summer where biomass burning migrates from subequatorial Africa to the northern tropics but where interannual variations such as El-Niño Southern Oscillation (ENSO) can have a significant impact on burning over Indonesia and Australia (Thompson et al., 2001).

Biomass burning will generally produce a number of hydrocarbons for which carbon monox-

ide is an important tracer. Observations of CO vertical profiles from TES are used to examine the distribution of pollution generated from biomass burning in the southern Hemisphere. The key chemical mechanism for ozone production involves the NO_x ($\text{NO} + \text{NO}_2$) family. Observations of NO_2 tropospheric columns from the Ozone Monitoring Instrument (OMI) are used to show regions of enhanced surface emissions. Colocation of enhanced values of both CO and NO_x provides the critical ingredients for anthropogenic ozone formation. Complicating this analysis, however, is the production of NO from lightning, which is particularly intense over the tropics (Hauglustaine et al., 2001; Martin et al., 2002; Sauvage et al., 2007). Observations of lightning flash counts from the Lightning Imaging Sensor (LIS) are used to get a sense of the zonal distribution of lightning and its role in ozone formation.

Tropical tropospheric ozone has been studied extensively from a variety of platforms including aircraft (Marenco et al., 1998), ships (Thompson et al., 2000), sondes (Logan and Kirchoff, 1986; Thompson et al., 2003b; Oltmans et al., 2001), and satellites (Fishman et al., 1991). Of these observations, the Southern Hemispheric Additional Ozonesondes (SHADOZ) network of ozone sonde observations has provided the longest and most extensive record of the vertical distribution of ozone. Ozone measured from this network for November 2004 provides important correlative information.

These datasets provide the observational context to relate surface emissions, ozone precursors, ozone, and the pollution pathways connecting them. We quantify this relationship through the GEOS-Chem chemistry and transport model and optimal linear parameter estimates of surface emissions. Carbon monoxide is a good proxy for combustion byproducts. (Jones et al., 2007) conducted an inverse analysis of CO emissions for November 2004 using TES and Measurements Of Pollution In The Troposphere (MOPITT) data as constraints. The model was then run using the *a posteriori* CO emissions, and with changes in NO_x and hydrocarbon emissions scaled to the changes in CO emissions, to give updated ozone fields. Here we compare these ozone fields with TES observations of ozone. Residual differences between TES and model ozone are investigated by analyzing the differences in ozone, CO, NO_x , and PAN between *a priori* and *a posteriori* emissions.

2 Tropospheric Emission Spectrometer

2.1 Introduction

TES is an infrared, high resolution, Fourier Transform spectrometer covering the spectral range $650 - 3050 \text{ cm}^{-1}$ (3.3 - 15.4 μm) at a apodized spectral resolution of 0.1 cm^{-1} (nadir viewing). Launched into a polar sun-synchronous orbit (13:38 hrs local mean solar time ascending node) on July 15, 2004, the TES orbit repeats its ground track every 16 days, allowing global mapping of the vertical distribution of tropospheric ozone and carbon monoxide along with atmospheric temperature, water vapor, surface properties (nadir), and effective cloud properties (nadir). TES has a fixed array of 16 detectors, which in the nadir mode, have an individual footprint of approximately $5.3 \times .5 \text{ km}^2$. In order to increase the signal-to-noise ratio, these detectors are averaged together to produce a combined footprint of $5.3 \times 8.4 \text{ km}^2$. TES has two basic observational modes: the global survey mode, where observations are taken 5 degrees apart in latitude, and the “step-and-stare” mode, where the separation between observations is approximately 40 km along the orbit (Beer and Glavich, 1989; Osterman and et al, 2007). For this study, 6 global surveys over the course of 12 days were used where each global survey mode produced 1152 observations per day. The data used here is based on V002, which is available at the NASA Langley Atmospheric Data Center (<http://eosweb.larc.nasa.gov/>). Both TES CO and ozone profile estimates have been compared against a variety of aircraft, in-situ, and model studies. TES ozone is biased high, particularly in the upper troposphere, by 3-10 ppb, compared to sondes (Nassar et al., 2007; Osterman et al., 2007; Worden et al., 2007) and lidar (Richards et al., 2007). TES CO profiles are within 15% of aircraft profiles (Luo et al., 2007a; Lopez et al., 2007) while TES CO columns are within 4.4% of MOPITT columns (Luo et al., 2007b).

2.2 Characterization of TES trace gas profile estimates

The estimate of an atmospheric state, e.g., vertical distribution of ozone, is calculated through the minimization of the norm difference between spectral radiances measured by TES and an atmospheric “forward” model subject to constraints on the first and second-order statistics of

that atmospheric state. This minimization is carried out through a non-linear least squares optimization algorithm. A detailed linear error analysis is performed around the estimated state that accounts for random, systematic, and “smoothing” and “cross-state” error (Bowman et al., 2006, 2002; Worden et al., 2004)

Under the assumption that difference between the estimated and true state is linear with respect to the difference in spectral radiances, the estimated state can be related to the true state through the following linear model:

$$\hat{\mathbf{x}} = \mathbf{x}_a + \mathbf{A}(\mathbf{x} - \mathbf{x}_a) + \epsilon \quad (1)$$

where $\hat{\mathbf{x}}$, \mathbf{x} , \mathbf{x}_a are the estimated, “true”, and *a priori* state vectors, respectively, ϵ is the observational error with covariance

$$\mathbf{S}_\epsilon = E[\epsilon\epsilon^\top] \quad (2)$$

that accounts for the random, systematic and “cross-state” error terms (Worden et al., 2004). The averaging kernel matrix, \mathbf{A} , can be defined as

$$\mathbf{A} = \frac{\partial \hat{\mathbf{x}}}{\partial \mathbf{x}}. \quad (3)$$

The averaging kernel matrix defines the sensitivity of the estimated state to changes to the true state. The averaging kernel matrix is used to calculate the vertical resolution, information content, and degrees of freedom for signal of the estimate or “retrieval” (Rodgers, 2000). The averaging kernel is a non-linear function of forward model parameters, e.g. cloud optical depth, as well as the retrieved state. For example, higher ozone concentrations result in greater sensitivity and therefore higher values in the averaging kernel. Figure 1 shows the average of the diagonal of the averaging kernel matrix from 15S to the equator as a function of longitude for TES estimates of ozone for the November 4-16 time period. Larger values indicate greater sensitivity to the atmospheric state at their corresponding pressure levels. The peaks of the averaging kernel matrix are centered near 600 mb indicating that TES observations have significant sensitivity to the lower troposphere.

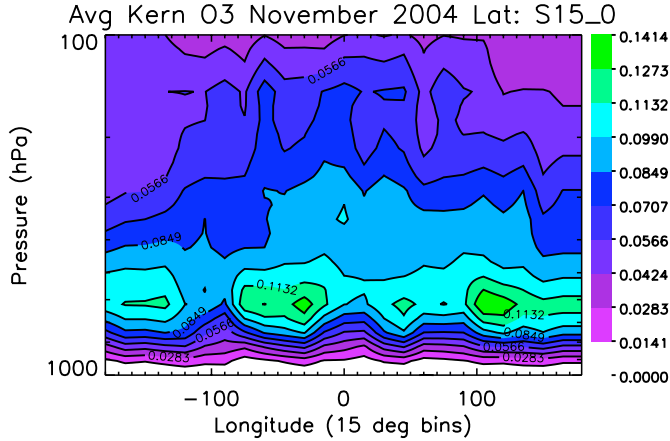


Fig. 1. Mean of the ozone averaging kernel diagonals for TES observations from 15S to the equator. The mean values are calculated in 15 x15 degree bins.

A suite of quality criteria are used for selection of the observations. For ozone, the absolute radiance residual means is less than 0.1, the radiance root mean square values are between 0.5 and 1.75, the retrieved cloud top pressure is between 90 and 1300 hPa, the absolute difference between surface temperature and atmospheric temperature is less than 25K, the absolute difference of the emissivity from its *a priori* value is less than 0.04, and the absolute difference between the surface temperature and its *a priori* value is less than 8K (Osterman and et al, 2007).

2.3 Construction of the TES observation operator and comparison to chemistry and transport models

The vertical resolution and bias characterized by Equation (3) must taken into account in order to compare TES ozone and CO profile estimates with *in-situ* measurements and modeled profiles. The TES observation operator is constructed to perform this function and will be shown for comparison with a chemistry and transport model (CTM). A CTM can be described by a “forward” model

$$\mathbf{x}_t^{i,m} = \ln \mathbf{F}^i(\mathbf{y}_t, \mathbf{u}_t, t) \quad (4)$$

where $\mathbf{x}_t^{i,m}$ is the vector whose elements are the natural logarithm of the vertical distribution of the model atmospheric state, e.g., CO, at location i and time t , \mathbf{y}_t is a vector whose elements are the 3-D distribution of the atmospheric state, \mathbf{u}_t is a vector whose elements contain key source and sink terms for the atmospheric state, and \mathbf{F} is the model operator that interpolates the global atmospheric state to the atmospheric state at location i to the TES footprint. The TES observation operator is

$$\mathbf{H}_t^i(\mathbf{x}_t, \mathbf{u}_t, t) = \mathbf{x}_{t,a}^i + \mathbf{A}_t^i(\mathbf{x}_t^{i,m} - \mathbf{x}_{t,a}^i). \quad (5)$$

The natural logarithm operation on the CTM model operator in Equation (4) accounts for the fact that TES retrievals of trace gases such as ozone and CO are performed on the natural logarithm of those gases. By implication, the *a priori* state vector and averaging kernel matrix are also in natural logarithm and consequently the statistics are assumed to be lognormal in distribution. In the case where the actual atmospheric state is equal to Equation (4), then the TES profile estimate can be written in the standard noise model

$$\hat{\mathbf{x}}_t^{i,m} = \mathbf{H}_t^i(\mathbf{y}_t, \mathbf{u}_t, t) + \epsilon. \quad (6)$$

Equation (6) includes both the vertical resolution and characterized errors in the TES retrieval. Subtracting Equation (5) from (1) results in

$$\hat{\mathbf{x}}_t^i - \hat{\mathbf{x}}_t^{i,m} = \mathbf{A}_t^i(\mathbf{x} - \mathbf{x}_t^{i,m}) + \epsilon. \quad (7)$$

where the averaging kernel varies as a function of location and time. The bias associated with the *a priori* is removed in the comparison between the model and the TES retrieval in Equation (7). The first term on the right hand side of Equation (7) accounts for the vertical resolution of the estimate and the second term accounts for the observational error. This approach was used to demonstrate the potential of TES observations to constrain CO emissions in (Jones et al., 2003).

3 Overview of TES tropical tropospheric ozone and carbon monoxide observations

TES observations of ozone and CO are shown from 15N to 30S at 464 hPa from November 4-16, 2004 in Figure (2). The most notable feature is a band of elevated ozone starting from eastern Brazil through both the Atlantic and Indian Oceans and extending into the Pacific. The highest ozone concentrations are observed both over the tropical Atlantic (>100 ppbv) and over Madagascar. This pervasive zonal ozone distribution has been observed from satellites, in particular from the total ozone mapping spectrometer (TOMS) using a tropospheric ozone residual technique (Fishman and Larsen, 1987; Fishman et al., 1991, 2003). This distribution is due in part to the recirculation of ozone and ozone precursors between South America and sub-equatorial Africa over the Atlantic (Kalnay et al., 1996; Krishnamurti et al., 1996; Thompson et al., 1996; Sinha et al., 2004).

In addition, a high pressure system centered over Australia, low averaged cloud optical depths over November from the International Satellite Cloud Climatology Project (ISCCP) (Rossow and Schiffer, 1991; Rossow et al., 1993) (available at <http://isccp.giss.nasa.gov/>), and relatively high biomass burning (van der Werf et al., 2006) indicate conditions favorable to ozone formation. TES observations of mid-tropospheric ozone show enhanced values extending northwest of Australia into Indonesia, which have been associated with El Niño conditions (Thompson et al., 2001; Chandra et al., 2007).

TES observations of CO show a plume of CO from South America and extending into the

western Pacific consistent with previous satellite and aircraft observations (Chatfield et al., 2002; Edwards et al., 2006). These concentrations in conjunction with MODIS firecounts are indicative of a continued presence of continental emission sources even as the southern hemisphere transitions to its Austral summer, wet season. Similar to TES ozone, Indonesia-Australia region shows elevated concentrations of TES CO comparable to South America and sub-equatorial Africa. In addition, the pervasive high values of CO across the Indian ocean are suggestive of recirculation of emissions between continents, which is consistent with studies from the Southern African Fire-Atmosphere Research Initiative (SAFARI), e.g., (Garstang et al., 1996).

3.1 Comparison of TES ozone to the SHADOZ network

The vertical distribution of ozone over the southern tropics as observed by TES is shown in Figure 3 where the TES observations have been averaged in 15° bins between the equator and 30S. There were roughly 30 observations for each bin. A pervasive high in ozone is evident across the tropical Atlantic with values up to about 80 ppbv from 15S to the equator. This distribution follows the so-called "wave-one" pattern (Thompson et al., 2000; Logan, 1999). From Figure 3(a) there is a secondary ozone enhancement over Indonesia-northern Australia between 90E-100E and 400-500 hPa. A similar picture emerges based on ozone sondes drawn from the SHADOZ network (Thompson et al., 2003a) for November 2004, which is shown in Figure(4). A total of 30 sondes were used in the average ranging from just one sonde measurement at Java to 6 sonde measurements at Natal. In the tropical Atlantic between 0 and 30W, Ascension (8S, 14.4W) and Natal (5.8S, 35.2W) show middle tropospheric values between 60-80 ppb, consistent with TES observations in Figure 3. At the Java site (7.5S, 112.6E), elevated ozone concentrations of 50-70 ppb are observed between 700-400 hPa while TES observations over the same region indicate a similar enhancement. The vertical structure of the ozone over Indonesia is somewhat different than ozone enhancements over the tropical Atlantic and western Indian Ocean suggesting that different processes are controlling ozone formation there.

The vertical distribution of TES ozone estimates from 30S to 15S are shown in Figure 3(b). Elevated ozone stretches from Southern Brazil across the Atlantic and Africa into most of the In-

dian Ocean. This elevated ozone is pervasive from roughly 500-200 hPa. Comparison between Pretoria (25.9S, 28.2E) and TES observations show similar values of ozone (80-100 ppb) between 400-200 hPa whereas Reunion Island (21.1S, 55.5E) indicates significantly higher ozone above 200 hPa. Similar to the ozone distribution in Figure 2, higher amounts of ozone are seen throughout the troposphere over the Indian Ocean relative to the remote Pacific by roughly 10-20 ppb, consistent with transport of ozone from South America, South Atlantic, and Africa into the Indian Ocean.

4 Signatures of lightning and surface NO_x

The concentrations and distribution of NO_x has a significant impact of the distribution of ozone (Jacob et al., 1996). In the southern hemisphere, the primary sources of surface NO_x are biomass burning, fossil fuel and biofuel combustion (Jaeglé et al., 2005). These emissions can produce ozone near the surface which can in turn be convectively lofted into the upper troposphere (Chatfield and Delany, 1990). However, NO_x from lightning is directly emitted into the upper troposphere and can play a dominant role in the production of tropical ozone (Pickering et al., 1998; Sauvage et al., 2007; Martin et al., 2007; Boersma et al., 2005). The Lightning Imaging Sensor (LIS) aboard the Tropical Rainfall Measuring Mission (TRMM) estimates lightning flash counts by means of a high speed CCD imaging sensor (3-6 km horizontal resolution) in conjunction with a narrow band ($\lambda = 777\text{nm}$) filter. Lightning flash counts from LIS are shown in Figure 5 for November 2004. For this month, lightning flash counts are densely distributed over Northern Argentina and to a lesser extent southeastern Brazil, throughout tropical Africa and Southern Africa with rates exceeding 150. By comparison, Indonesia/Northern Australia shows markedly less flash counts with rates generally less than 25. This distribution is consistent with the high pressure system from the NCEP reanalysis and the ISCCP cloud optical depth. Consequently, we could expect the regional contribution of ozone from lightning NO_x over Indonesia/Australia to be less than the regional contribution of lightning to South America and Africa.

The distribution of lower tropospheric NO₂ can be investigated from monthly average tro-

pospheric NO_2 columns derived from the Ozone Monitoring Instrument (OMI, (Levelt et al., 2006)), which are shown in Figure 6 for November 2004. The columns are calculated using the retrieval-assimilation algorithm described in (Boersma et al., 2004, 2007). Individual OMI tropospheric NO_2 observations with approximate horizontal resolutions of $25 \times 24 \text{ km}^2$ have been gridded onto a $0.5^\circ \times 0.5^\circ$ grid. To avoid situations with clouds screening the NO_2 underneath, only cloud-free (cloud radiance fraction $< 50\%$) observations were taken. The estimated uncertainty for individual OMI observations is on the order of 30-50% for situations with appreciable NO_2 columns ($> 1 \times 10^{15} \text{ molec/cm}^2$), but it is anticipated that the averaging of large numbers of pixels here reduces the uncertainty of the monthly average to within 5-10%. Given these uncertainties, NO_2 tropospheric column values on the order of $8 \times 10^{15} \text{ molec/cm}^2$ are concentrated south of the mouths of the Amazon in Brazil as well as Northern Australia. With the exception of Johannesburg region in South Africa where values approach $20 \times 10^{15} \text{ molec/cm}^2$, there are no high concentrations of tropospheric NO_2 in sub-equatorial Africa.

5 Comparison of GEOS-Chem to TES estimates of CO and ozone

5.1 Description of GEOS-Chem

The GEOS-Chem global chemistry and transport model was originally described by (Bey et al., 2001). The simulation conducted for the November 2004 used GEOS-Chem v7.02.04 (<http://www-as.harvard.edu/chemistry/trop/geos>) driven by GEOS-4 assimilated meteorological observations from the NASA Global Modeling and Assimilation Office (GMAO). The GEOS-4 observations have a temporal resolution of 6 hours (3 hours for surface variables and mixing depths), a horizontal resolution of $1^\circ \times 1.25^\circ$, and 55 vertical layers. Here we degrade the horizontal resolution to $2^\circ \times 2.5^\circ$ from the surface up to 0.01 hPa. The model includes a complete description of tropospheric O_3 - NO_x -hydrocarbon chemistry, including sulfate aerosols, black carbon, organic carbon, sea salt, and dust. Anthropogenic emissions in the model are described in (Duncan et al., 2007). Extensive evaluations of the GEOS-Chem tropospheric ozone simulations have been conducted by (Wu et al., 2007; Martin et al., 2002; Liu et al., 2006).

5.2 Comparison of GEOS-Chem to TES CO over the southern tropics

The zonal distribution CO from the GEOS-Chem model from 15°S to the equator is shown in Figure 7(a). The results are averaged from November 4-16, 2004 in 15° × 15° bins. This simulation used the climatological biomass burning, which gives elevated values of CO over South America, Africa, and Indonesia/Australia. In the lower and middle troposphere CO over South America dominates the region with values up to 40 ppb higher relative to Indonesia/Australia at the surface. The zonal distribution of TES CO estimates from 15°S-0 is shown in Figure 8(c). These retrievals are also averaged in 15° longitudinal bins with roughly 20-30 observations per bin. For comparison, the GEOS-Chem CO fields were sampled at the coincident TES observation coordinates and the TES observation operator, (Equation 5) was applied as shown in Figure 8(a). There is significant disagreement both in the magnitude and relative distribution of the GEOS-Chem and TES CO observations with differences up to 40 ppb. TES observations in Figure 8(c) show that CO over Indonesia/Australia was as high as that over South America.

TES CO observations were used to estimate the CO source emissions over the globe in (Jones et al., 2007). The *a priori* and *a posteriori* emissions for South America, sub-equatorial Africa, and Australia/Indonesia are listed in Table 1. For this time period, the emissions were estimated to be over twice as high as those in the *a priori* simulation. The GEOS-Chem results at the TES resolution and sampling with the *a posteriori* emission are shown in Figure 8(b). The *a posteriori* CO field distribution from GEOS-Chem between 15°S and the equator is in remarkably good agreement with the TES observations shown in Figure 8(c).

The response of GEOS-Chem CO fields to changes in the emissions is shown in Figure 7(b). The maximum increase in CO is over the Indonesia/Australia region is almost 100 ppb or 85% near the surface and approximately 60 ppb or a 65% increase throughout the free troposphere. Over the Indian Ocean, the CO distribution in GEOS-Chem increased by about 35 ppb over the Indian Ocean and around 45 ppb over sub-equatorial Africa in the 200-400 hPa region. Over South America, the increase is modest - no more than 30 ppb.

5.3 Comparison of GEOS-Chem to TES ozone over the southern tropics

The zonal distribution of ozone from the GEOS-Chem model with *a priori* emissions is shown in the top panel of Figure 9(a) from 15°S-0 averaged in the same manner as CO. GEOS-Chem follows the familiar “wave-one” pattern (Thompson et al., 2003b) with enhanced values of ozone across the tropical Atlantic. However, there is a modest secondary maximum in ozone over Indonesia/Australia relative to the Pacific. This enhancement is also observed in Figure 10(a) where the TES observation operator has been applied to the GEOS-Chem fields. In both case, the ozone amounts are less than those observed by TES in Figure 3 over both the tropical Atlantic and Indonesia/Australia.

The ozone distribution from GEOS-Chem was also calculated based on the revised emissions where the NO_x emissions were scaled with the CO *a posteriori* emission estimates. The GEOS-Chem fields with the *a posteriori* emissions sampled along the TES observations are shown in Figure 10(b). There is an increase in upper tropospheric ozone at 200 hPa over the tropical Atlantic and at 280 hPa over sub-equatorial Africa. In addition, an overall increase of about 10 ppb throughout the troposphere can be seen over Indonesia/Australia. Use of the *a posteriori* emissions improves agreement between the model and TES ozone, but significant discrepancies remain.

The difference between the TES observations of ozone and GEOS-Chem with *a priori* (top panels) and *a posteriori* (bottom panels) emissions are shown in Figure 11 for 15S to the equator and Figure 12 for 30S to 15S. The top panels show the largest differences in ozone are centered over the Atlantic as well as Indonesia. With the *a posteriori* emissions, the bottom panels show an overall decrease in ozone differences that is fairly uniform zonally. Over the tropical Atlantic, the difference between GEOS-Chem and TES are reduced by roughly 5 ppb from 30S-0. From 30S-0 over the Indonesia/Australia region in the mid-troposphere the reduction is more substantial with reductions up to 10 ppb. On the other hand, the upper tropospheric ozone difference at 100E and 100W from 15S-0 has increased with the *a posteriori* emissions. With those exceptions, TES observations are higher everywhere relative to GEOS-Chem.

In contrast to the *a posteriori* CO comparisons with TES observations (Figure 8), there re-

Table 1. A priori and *a posteriori* emissions taken from (Jones et al., 2007).

Region	a priori (Tg CO/y)	a posteriori
S. America	113	118
S. Africa	95	173
Indonesia/Australia	69	155

mains significant residual differences in ozone that have a spatial structure. This residual indicates that assumptions used in the emissions are incorrect, e.g., the relative distribution of CO to NO_x, pathways relating those emissions to ozone formation are deficient, or background processes have not been properly described. Over the Atlantic, the residual differences and their spatial structure can be attributed in part to ozone generated from lightning NO_x. In (Sauvage et al., 2007), increasing the intra-cloud to ground-to-cloud flash ratio to 0.75 for lightning NO_x formation considerably improved agreement between GEOS-Chem and SHADOZ network ozone for the September-October-November season (although this increase reduced agreement in other seasons). The peak changes in ozone to this ratio were centered between 500-300hPa over the Ascension Islands and increased ozone there by 10-20 ppb, which is consistent with the residual difference in Figure 11(b). In addition, there is a residual difference over Indonesia/Australia at 600 hPa of up to 15 ppb that can not be explained by surface emissions. This difference may reflect deficiencies in sources of NO_x from lightning in the region, vertical mixing, or assumed composition of the emission sources.

5.4 Response of GEOS-Chem to changes in ozone and NO_x distribution from *a posteriori* emission estimates

We can use the emission estimates to investigate chemical mechanisms linking those emissions to the tropical ozone distribution and to interpret the residual differences between TES and GEOS-Chem ozone distributions. The averaged difference between GEOS-Chem ozone fields with *a priori* and *a posteriori* emissions are shown in Figure 9. The largest differences in ozone from the change in emissions are over the Indonesia/Australia regions where ozone increases by

up to 16 ppb in the upper troposphere centered around 150 hPa. It is in this upper tropospheric region, as shown in Figure 11(b), that GEOS-Chem ozone is greater than the TES observations by up to 15%. The amount of ozone produced, however, will be sensitive to the chemical composition of the lofted emissions. Consequently, one interpretation is that the overestimate is a consequence of the uniform scaling of all combustion sources.

The ozone response to the emission changes over sub-equatorial Africa is approximately 8 ppb near the surface and around 200 ppb. Over South America, there were few changes in the ozone distribution, consistent with a modest increase in emission strengths. Curiously, there was a significant increase in ozone in the remote Pacific centered around 150S in the upper troposphere ($>15\%$).

The principle chemical mechanism for the ozone response in the free troposphere to changes in surface emissions is the ambient NO_x distribution. CO is assumed to be a tracer of emissions generally and consequently all the emissions, including NO_x , are scaled along with the CO emissions derived from the inverse analysis. However, the NO_x zonal distribution has a different response to the scaled emissions than the CO distribution. The NO_x distribution based on the GEOS-Chem *a priori* emissions and the change in mean zonal NO_x from the *a posteriori* emissions are shown in Figure 13. The *a priori* NO_x fields are highest over South America where the values are up to 6-7 times higher than over Indonesia/Australia and up to twice as high as sub-equatorial Africa. The concentrations of NO_x in the free troposphere are due primarily to lightning sources (Pickering et al., 1998; Folkins et al., 2006), with the South American and sub-equatorial African regions exhibiting a much larger source of NO_x from lightning than the Indonesian/Australian regions. Qualitatively, this distribution is consistent with the LIS observations in Figure 5. Associated with the higher concentrations of NO_x , the model simulation with the *a priori* emissions also produces more ozone (Figure 9) over South American and sub-equatorial Africa than over Indonesia/Australia. The low ozone abundance over Indonesia/Australia, however, also reflects convective transport of ozone-poor marine air to the upper troposphere (Lelieveld et al., 2001). Enhanced ozone over South America and sub-equatorial Africa results in greater concentrations of OH (by more than a factor of 2) over these regions, which together with the higher NO_x levels leads to significantly more HNO_3

(by almost an order of magnitude) over South American and sub-equatorial Africa relative to Indonesia/Australia.

The greatest increase in free tropospheric NO_x (100 ppt) to the *a posteriori* emissions is centered over the Java Sea (115E) at 150 hPa just to the east of the high NO_x concentrations over Sumatra (105E). Conversely the greatest decrease (>120 ppt) in free tropospheric NO_x is located over the western coast of Africa. In addition, there is a significant decrease over South America (>55 ppt) centered at 250 hPa. The response of free tropospheric NO_x to increases in the surface emissions, which include surface NO_x , is a non-linear function of both the ambient amounts of ozone, NO_x , and OH along with the chemical composition of lofted emissions. Over Indonesia/Australia the dominant sink for NO_x is formation of peroxyacetylnitrate (PAN), whereas over South America and Africa NO_x is lost through formation of PAN and HNO_3 (due to the higher levels OH in these regions). In addition, the NO/NO_2 ratio is lower over South America and Africa because of the higher abundances of ozone in these regions. This enhances the conversion of NO_x to PAN and HNO_3 .

Another important difference between the three tropical continental regions is the distribution of organics such as acetaldehyde, acetone, and formaldehyde in the free troposphere. Acetaldehyde, for example, is oxidized by reaction with OH to produce peroxyacetyl radicals ($\text{CH}_3\text{C}(\text{O})\text{OO}\cdot$) that in turn react with NO_2 to form PAN. The mean difference in PAN concentrations between the simulations with the *a priori* and *a posteriori* emissions is shown in Figure 14. The response of upper tropospheric PAN to surface emission changes above Africa is over 150 ppt, which is roughly 50% greater than over Indonesia/Australia. Tropospheric PAN also increased over South America with changes up to 100 ppt. We can conclude that, for this time period, increases in surface emissions over South America and sub-equatorial Africa preferentially lead to the formation of PAN at the expense of NO_x and consequently mute the production of ozone. On the other hand, increased surface emissions in Indonesia/Australia, while leading to enhanced PAN, do not lead to a reduction of NO_x due to the overall lower background concentrations of NO_x , OH, and carbonyl compounds. Consequently, ozone production is regionally enhanced. The different responses to increased emissions over these three regions illustrate the importance of both background meteorological conditions and the partic-

ular chemical composition of the emissions in linking ozone production to surface emissions. These responses must be characterized in order to reduce reduce uncertainty both in present day and future changes in ozone (Horowitz, 2006).

6 Conclusions

We have investigated the processes controlling the zonal distribution of tropical tropospheric ozone with a focus on the sensitivity of that distribution to changes in surface emissions between South America, sub-equatorial Africa, and Indonesia/Australia for November, 2004.

Against the backdrop of the “wave-one” pattern of elevated ozone in the tropical Atlantic, TES ozone profiles also indicate enhanced values over Indonesia/Australia with volume mixing ratios up to 70 ppb at 600 hPa. This enhancement is consistent with a SHADOZ sonde observation over Java. Co-located CO profiles from TES and NO₂ columns from OMI indicate concentrations over Indonesia/Australia are comparable to those over South America and Africa.

From this observational context, we assessed the contribution of surface emissions to tropical ozone using GEOS-Chem simulations with *a posteriori* emissions derived from a linear inverse model, which was based on TES and MOPITT CO, developed in (Jones et al., 2007). Based on over a factor of 2 increase in surface emissions in sub-equatorial Africa and Indonesia/Australia, the overall difference between TES and GEOS-Chem ozone was reduced throughout the troposphere between 30S-0. Over Africa and Indonesia/Australia the discrepancies between GEOS-Chem and TES decreased by roughly 10 ppb.

While there was overall improvement between TES ozone observations and GEOS-Chem, there remained substantial disagreements. Maximum residual differences of approximately 18 ppb are seen between 15S-0 and 30 ppb between 30S-15S. In the upper troposphere over the Eastern Indian Ocean and parts of the Western Pacific, GEOS-Chem overestimated the ozone distributions by 5 ppb.

The residual differences in ozone of 10-20 ppb in the mid-troposphere over the tropical Atlantic are consistent with the differences found in (Sauvage et al., 2007) associated with under-

estimates of lightning NO_x formation in GEOS-Chem for the September-October-November season. In addition, there is a residual difference over Indonesia/Australia at 600 hPa of up to 15 ppb that can not be explained by surface emissions.

We investigated these residual differences further by examining the spatial patterns in GEOS-Chem estimates of ozone, CO, and NO_x from changes between the *a priori* and *a posteriori* surface emissions. The greatest change to the free tropospheric ozone distribution from 15S-0 was over Indonesia (<16 ppb) at 175 hPa, consistent with maximum positive changes in NO_x (<100 ppt) and CO (<70ppb). Consequently, free tropospheric ozone over Indonesia/Australia is sensitive to changes in regional surface emissions and these emissions make a significant contribution to the regional ozone budget.

On the other hand, the free tropospheric NO_x distribution declined over Africa and South America with losses exceeding 150 ppt. We examined the PAN response as a possible loss mechanism for the NO_x . Maximum increases in PAN, which reached over 150 ppt, corresponded to the maximum decreases in the NO_x distribution. Therefore, conversion of NO_x to PAN can partially explain the decreases in NO_x in response to increases in surface emission over South America and Africa. If this mechanism is correct, then the sensitivity of the tropical Atlantic ozone to changes in surface emissions of NO_x is low because of the large ambient distribution of ozone and NO_x from lightning. However, the enhanced PAN could lead to additional ozone formation downwind through conversion of PAN back to NO_x (Staudt et al., 2003).

Based on scenarios discussed in the IPCC-4, the tropical latitudes are particularly sensitive to climate change in terms of precipitation and land-use (Solomon et al., 2007). Based on our results, the emissions from Indonesia/Australian are an important contributor to the zonal tropical ozone distribution both in terms of the ozone produced but also in the sensitivity of ozone to changes in those emissions. Given the complex feedbacks between land-use, biomass burning, biofuel production, plant productivity, and CO_2 uptake and emission, (Levine, 1999; Sitch et al., 2007; Lohman et al., 2007; Forster et al., 2007), quantifying the present and future impact of surface emissions to tropical ozone will be critical for understanding chemistry-climate coupling.

Acknowledgements. This work was performed, in part, at the Jet Propulsion Laboratory, California Institute of Technology, under a contract with the National Aeronautics and Space Administration (NASA). JAL was funded by a grant from NASA to Harvard University. DBAJ was supported by funding from the Natural Sciences and Engineering Research Council of Canada. We also thank the SHADOZ program for making the sonde data accessible.

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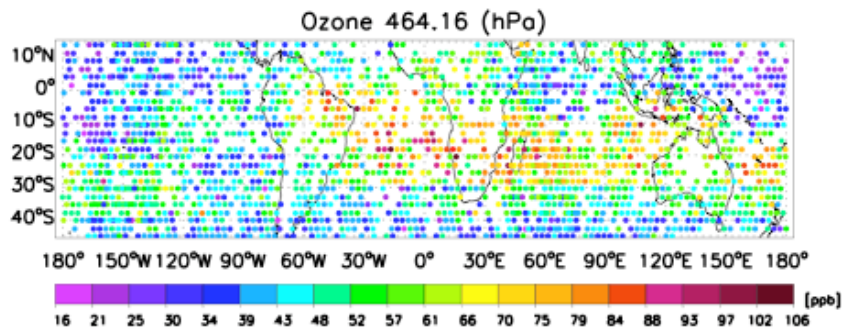
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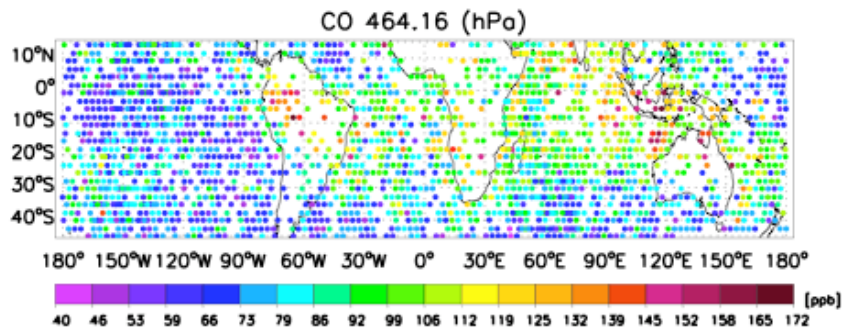
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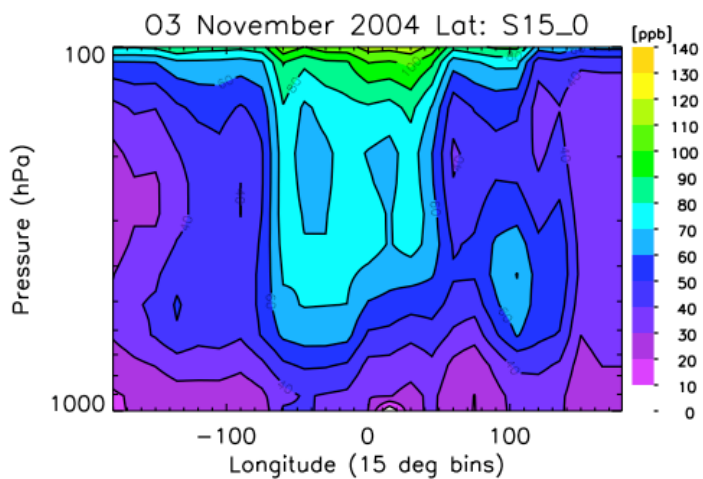


(a)

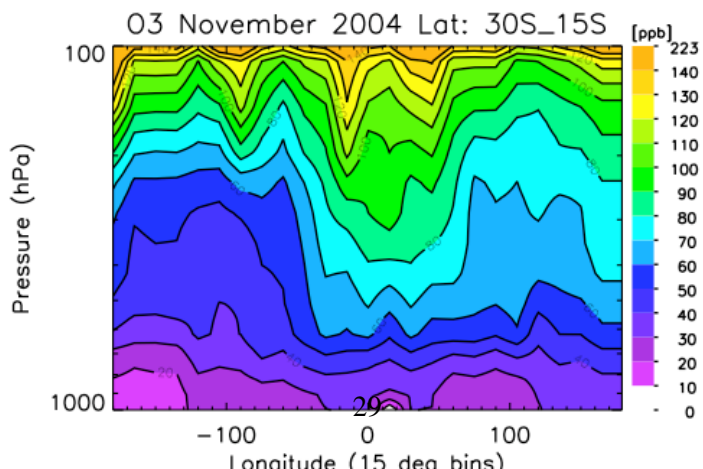


(b)

Fig. 2. (a) TES ozone estimates and (b) TES CO at 464.14 hPa from November 4-16, 2004 using V002 data.



(a)



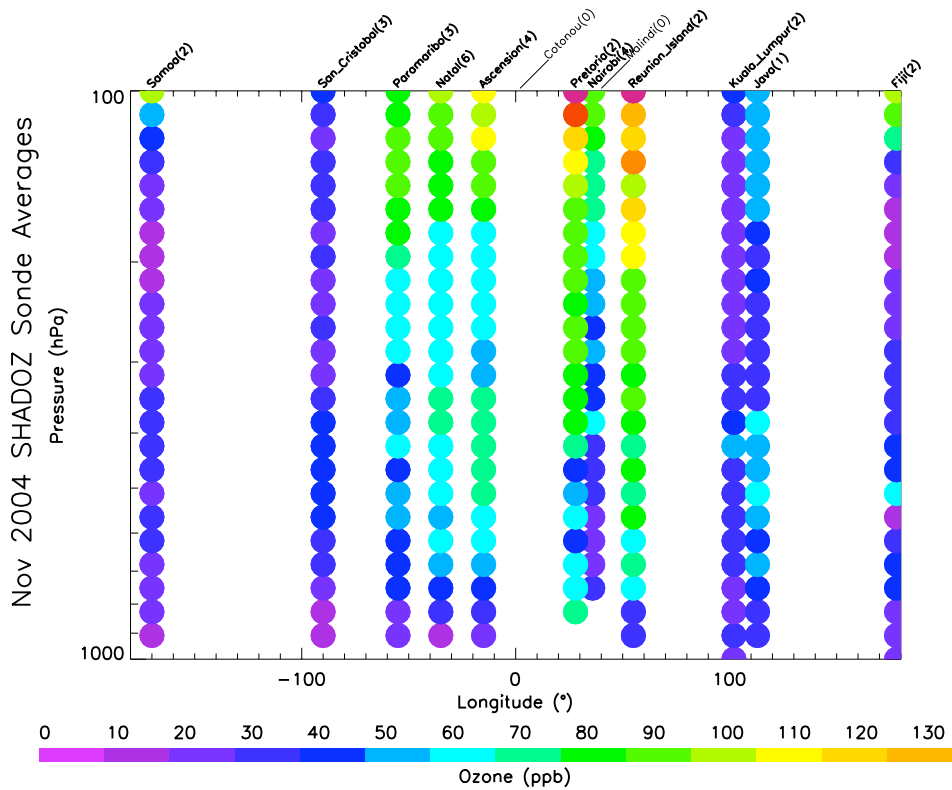


Fig. 4. SHADOZ ozone sonde measurements for November 2004. Location and the number of sondes used in the average are shown across the top of the figure. Most of the sites are between 0-15°S.

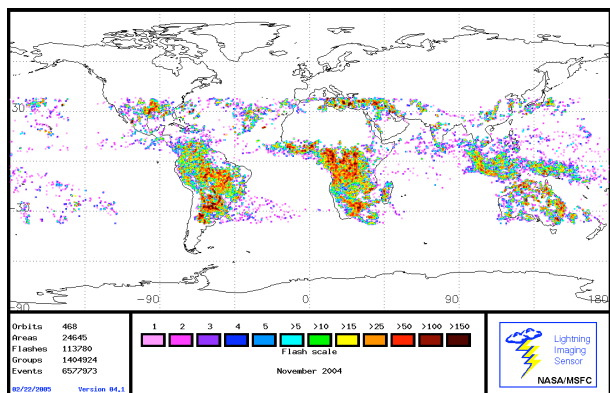


Fig. 5. Observations of lightning flash counts from the Lightning Imaging Sensor (LIS) for November 2004.

November 2004

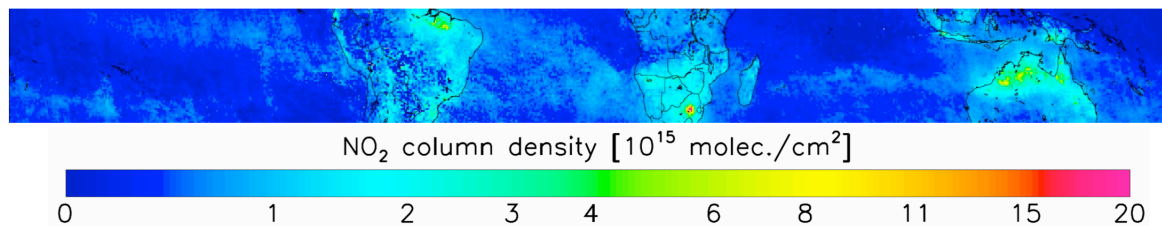
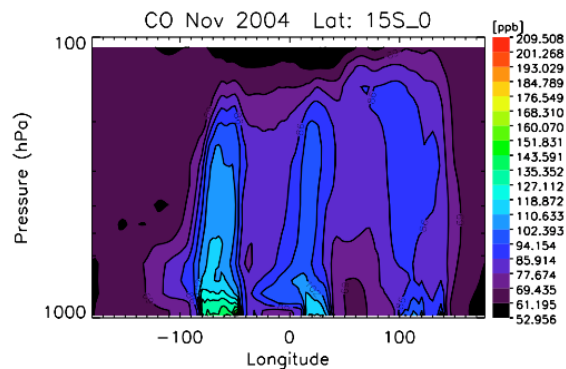
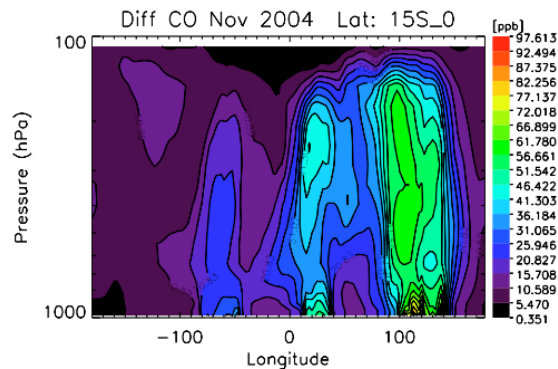


Fig. 6. OMI NO_2 tropospheric columns for November 2004.

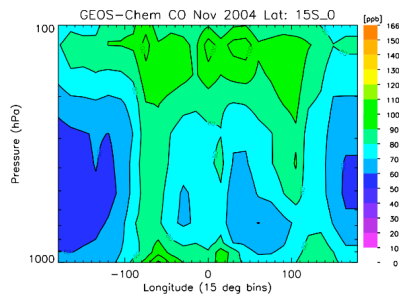


(a)

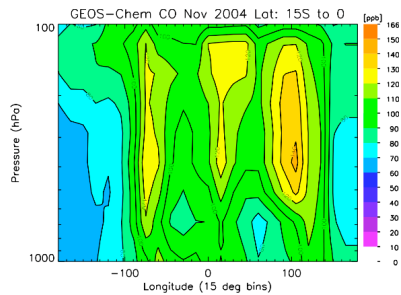


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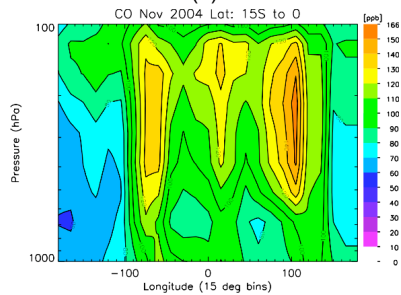
Fig. 7. (a) Zonal distribution of CO from GEOS-Chem from 15°S-0. The data is averaged in $15^\circ \times 15^\circ$ bins in both longitude and latitude. (b) Average difference in the GEOS-Chem zonal CO distribution between the *a priori* and *a posteriori* fields based on $15^\circ \times 15^\circ$ bins



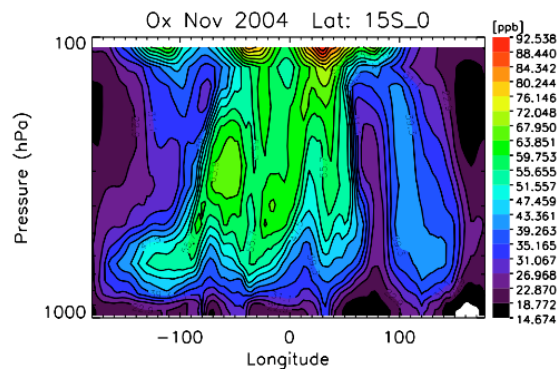
(a)



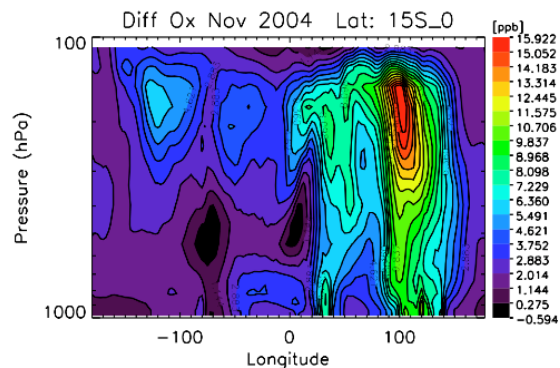
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(c)

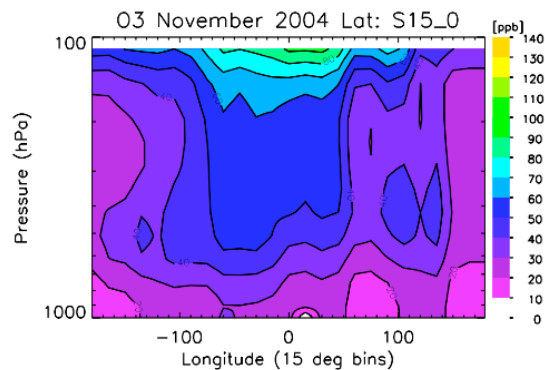


(a)

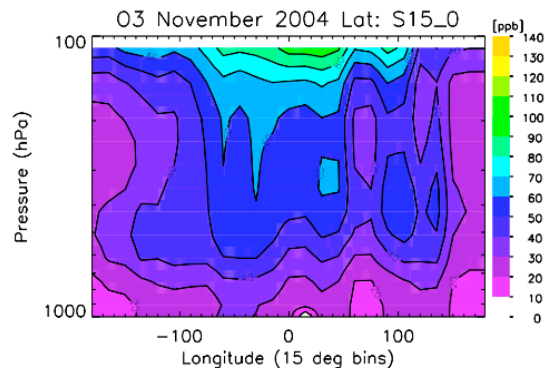


(b)

Fig. 9. (a) Zonal distribution of ozone from GEOS-Chem averaged in 15° bins in longitude and latitude from the equator to 15°S . (b) Average difference in GEOS-Chem ozone fields between *a priori* and *a posteriori* emissions based on $15^\circ \times 15^\circ$ bins



(a)



(b)

Fig. 10. (a) Zonal distribution of ozone from GEOS-Chem with the TES observation operator applied averaged from 15°S to the equator. The distribution is calculated from averaged 15° bins in longitude and latitude. (b) GEOS-Chem ozone fields with *a posteriori* emissions from the equator to 15°S sampled

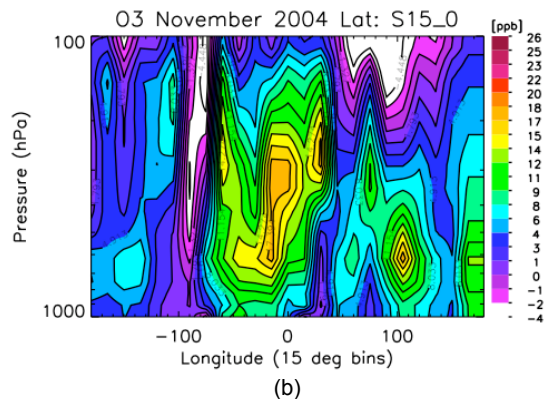
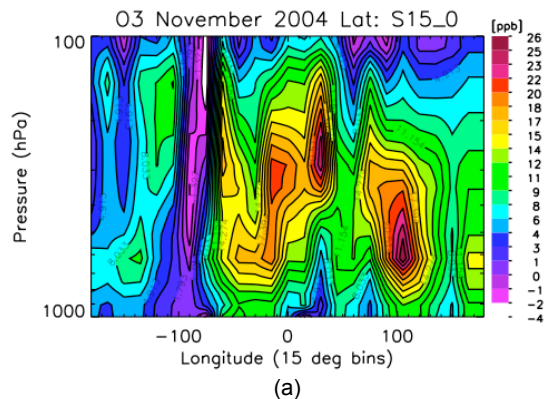
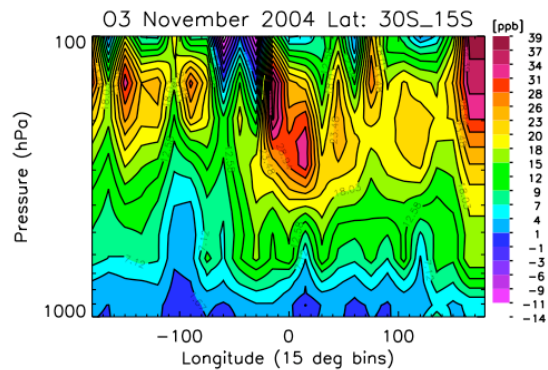
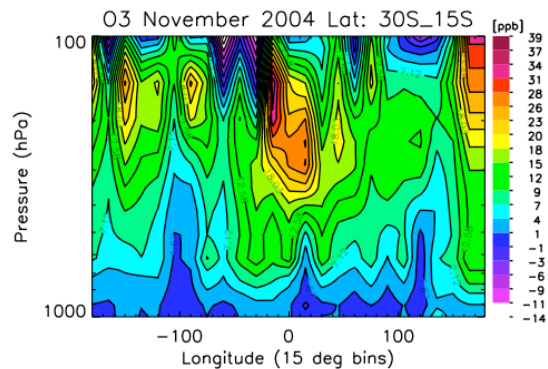


Fig. 11. (a) Mean difference between (a) GEOS-Chem with *a priori* emissions and TES ozone observations (b) GEOS-Chem with *a posteriori* emissions and TES ozone observations from 15S to the equator. Averages are calculated in $15^\circ \times 15^\circ$ bins

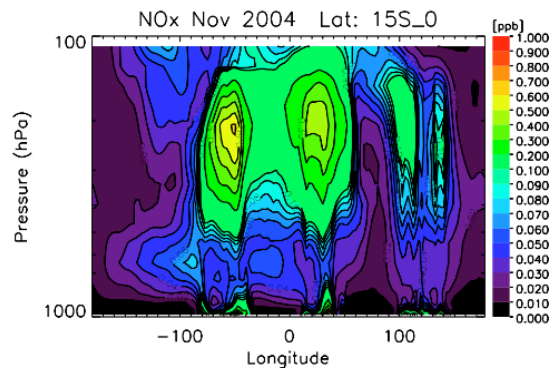


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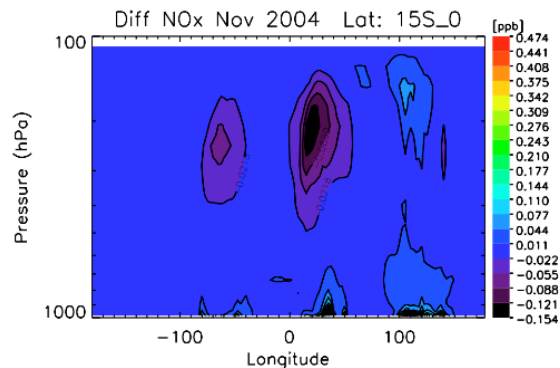


(b)

Fig. 12. Average difference between (a) GEOS-Chem with *a priori* emissions and TES ozone observations (b) GEOS-Chem with *a posteriori* emissions and TES ozone observations from 30S to 15S. Averages are calculated in $15^\circ \times 15^\circ$ bins.



(a)



(b)

Fig. 13. (a) Zonal NO_x concentrations in GEOS-Chem based on *a priori* emissions (b) Mean difference in the zonal NO_x distribution between *a priori* and *a posteriori* surface emission estimates between 15S and the equator during November 4-16 2004. Data is averaged over $15^\circ \times 15^\circ$ bins.

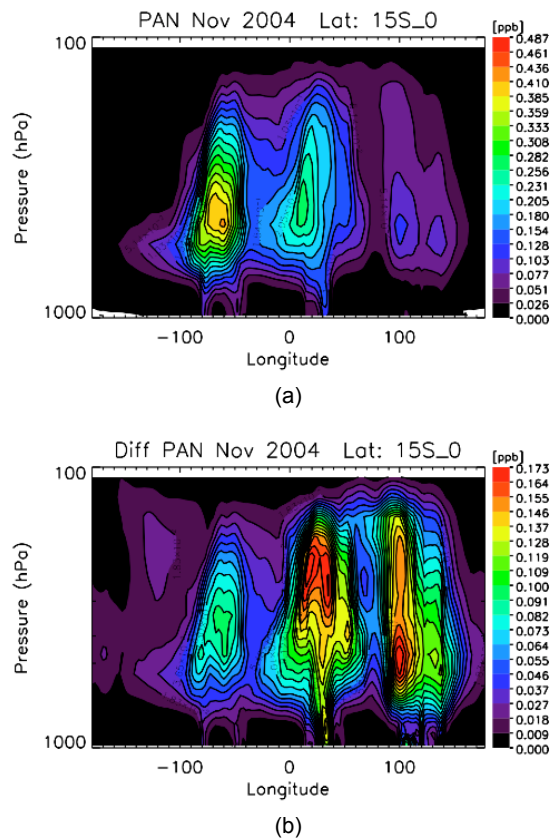


Fig. 14. (a) Zonal PAN distribution from GEOS-Chem. (b) Mean difference in the zonal PAN distribution between *a priori* and *a posteriori* surface emission estimates between 15S and the equator during November 4-16 2004. Data is averaged over $15^\circ \times 15^\circ$ bins.